```
L4 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2002 ACS
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AN 2001:923779 CAPLUS

DN 136:53771

TI Preparation of cyclic urea compounds

IN Rodriguez, Marc; Guichard, Gilles; Plaue, Serge; Semetey, Vincent; Schaffner, Arnaud-Pierre; Briand, Jean-Paul

PA Centre National de la Recherche Scientifique, Fr.; Neosystem; Galas-Rodriguez, Marie-Christine; Rodriguez, Pierre; Rodriguez, Elisa; Rodriguez, Romain

SO PCT Int. Appl., 103 pp.

CODEN: PIXXD2

DT Patent

LA French

FAN CNT 1

FAN.CNT 1				עדאים האשפ אחחו דכאשוראו אין האשפ													
PATENT NO.			K1.	KIND DATE				APPLICATION NO.					DATE				
ΡI	WO 2001096318			1	20011220			WO 2001-FR1837				7	20010613				
	W:	AE, A	5, AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
		CO, CI	R, CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	
		HR, H	J, ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,	
		LT, L	J, LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	
		RU, SI	, SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	
		VN, Y	J, ZA,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,	ТJ,	TM				
	RW:	GH, GI	1, KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	
		DE, D	K, ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	
		BJ, C	c, CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG			
	FR 2810039			A1 20011214				FR 2000-7507				20000613					
PRAI	PRAI FR 2000-7507				2000	0613											
OS GI	MARPAT	136:53	771														

The invention concerns a method for prepg. cyclic urea compds. from an activated carbamic acid deriv. contg. an unprotected primary or secondary amine function, by reaction between the primary or secondary amine function and the carbamic acid function of the carbamic acid deriv. Thus, the protected amine I was de-tert.-butoxycarbonylated and cyclized with EtN(CHMe2)2 to give the cyclic urea II.

IT 254100-96-4 254100-98-6

RL: RCT (Reactant); RACT (Reactant or reagent) (cyclization of amino carbamates to cyclic ureas)

RN 254100-96-4 CAPLUS

CN Carbamic acid, [(1S)-2-[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]-1-methylethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 254100-98-6 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-2-phenylethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RE.CNT 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
ANSWER 2 OF 6 CAPLUS COPYRIGHT 2002 ACS
L4
     2001:167650 CAPLUS
AN
    135:5262
DN
     (S)-O-Succinimidyl N-[2-(tert-butoxycarbonylamino)propyl]carbamate
TI
     Menschise, Valeria; Didierjean, Claude; Semetey, Vincent; Guichard,
AU
     Gilles; Briand, Jean Paul; Aubry, Andre
     Faculte des Sciences, Groupe Biocristallographie, UPRESA no 7036, Nancy
CS
I,
     Laboratoire de Cristallographie et Modelisation des Materiaux Mineraux,
et
     Biologiques (LCM3B), Universite Henri Poincare, Vandoeuvre les Nancy,
     54506, Fr.
     Acta Crystallographica, Section E: Structure Reports Online (2001),
SO
     E57(3), o222-o224
     CODEN: ACSEBH; ISSN: 1600-5368
     URL: http://journals.iucr.org/e/issues/2001/03/00/ya6006/ya6006.pdf
     International Union of Crystallography
PΒ
     Journal; (online computer file)
DT
     English
LΑ
     The mol. of activated carbamate, (S)-2, 5-dioxopyrrolidin-1-yl
AΒ
     N-[2-(tert-butoxycarbonylamino)propyl]carbamate,
     tBuOCONHCH(Me)CH2NHCOONC4H4O2 or Cl3H21N3O6, prepd. from
     N\textsubscript{-Boc-.beta.3HAla-OH,} assumes a folded conformation with the N\textsubscript{-C-N}
     torsion angle equal to 55.9 (3).degree.. Both N-H groups are involved
in
     intermol. hydrogen bonds, forming infinite chains in the crystal.
     254100-96-4
IT
     RL: PRP (Properties)
        (crystal structure; crystal structure of (S)-O-succinimidyl
        N-[2-(tert-butoxycarbonylamino)propyl]carbamate)
     254100-96-4 CAPLUS
RN
     Carbamic acid, [(1S)-2-[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]-
CN
```

Absolute stereochemistry. Rotation (-).

1-

RE.CNT 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

methylethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

App 5

```
ANSWER 3 OF 6 CAPLUS COPYRIGHT 2002 ACS
L4
AN
     2000:493513 CAPLUS
     133:105350
DN
     Preparation of stable activated peptide carbamic acids via azidolysis
\mathtt{TI}
and
     carbamovlation and use for preparing urea
     Rodriguez, Marc; Guichard, Gilles; Semetey, Vincent; Briand, Jean-Paul
IN
    Centre National de la Recherche Scientifique, Fr.; Galas-Rodriguez,
PA
     Marie-Christine; Rodriguez, Pierre; Rodriguez, Elisa; Rodriguez, Romain;
     Neosystem
     PCT Int. Appl., 174 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     French
LA
FAN.CNT 1
                                           APPLICATION NO. DATE
                            DATE
     PATENT NO.
                      KIND
                                                             20000114
                            20000720
                                           WO 2000-FR80
     WO 2000042009
                       A1
PI
         W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,
             CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,
             IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA,
             MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI,
             SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
             DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
             CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                             19990114
                                           FR 1999-330
                       Α1
                            20000721
     FR 2788518
                                            EP 2000-900588
                                                             20000114
     EP 1140822
                            20011010
                       A1
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
                            19990114
PRAI FR 1999-330
                       Α
                            20000114
                       W
     WO 2000-FR80
     CASREACT 133:105350; MARPAT 133:105350
OS
     The invention concerns the use of isocyanates obtained from amino acid
AB
     derivs. for prepg. and optionally isolating stable activated carbamic
acid
     peptides. or stable activated carbamates. Thus, Boc-Gly-gIle-CO2Su (Su
=
     succinimidyl) was prepd. from protected peptide Boc-Gly-Ile-OH in 4
steps
     via azidolysis and isocyanate intermediate with 87 % yield.
     284049-06-5
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. of stable activated peptide carbamic acids from protected
        peptides via azidolysis and carbamoylation reactions)
     284049-06-5 CAPLUS
RN
     Carbamic acid, [2-[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]-1-
CN
     methylethyl]methyl-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)
      O_C_NH_CH2_CH_Me
```

#### IT 254100-96-4P 254100-98-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. of stable activated peptide carbamic acids from protected peptides via azidolysis and carbamoylation reactions)

RN 254100-96-4 CAPLUS

CN Carbamic acid, [(1S)-2-[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]-1-methylethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 254100-98-6 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-2-phenylethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

IT 254100-97-5P 254100-99-7P 254101-00-3P

270575-71-8P 270575-72-9P 270575-73-0P 270575-74-1P 270575-75-2P 270575-76-3P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of stable activated peptide carbamic acids from protected peptides via azidolysis and carbamoylation reactions)

RN 254100-97-5 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-2-methylpropyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 254100-99-7 CAPLUS

CN Butanoic acid, 3-[[(1,1-dimethylethoxy)carbonyl]amino]-4-[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]-, phenylmethyl ester, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 254101-00-3 CAPLUS

CN Carbamic acid, [(1R,2R)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-2-(phenylmethoxy)propyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RN 270575-71-8 CAPLUS

CN Carbamic acid, [(1S)-2-[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]-1-methylethyl]-, 9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 270575-72-9 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-2-methylpropyl]-, 9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RN 270575-73-0 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-3-methylbutyl]-, 9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 270575-74-1 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-2-phenylethyl]-, 9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 270575-75-2 CAPLUS

CN Carbamic acid, [(1S)-2-[4-(1,1-dimethylethoxy)phenyl]-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]ethyl]-, 9H-fluoren-9-ylmethyl

ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 270575-76-3 CAPLUS

CN Carbamic acid, [(1S)-5-[[(1,1-dimethylethoxy)carbonyl]amino]-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]pentyl]-,
9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2002 ACS

AN 2000:177115 CAPLUS

DN 133:4952

TI Solid phase synthesis of oligoureas using O-succinimidyl (9H-fluoren-9-ylmethoxycarbonylamino)ethylcarbamate derivatives as activated monomers

AU Guichard, Gilles; Semetey, Vincent; Rodriguez, Marc; Briand, Jean-Paul

CS Laboratoire de Chimie Immunologique, UPR 9021 CNRS, Laboratoire de Chimie Immunologique, UPR 9021 CNRS, Institut de Biologie Moleculaire et Cellulaire, Strasbourg, 67084, Fr.

SO Tetrahedron Letters (2000), 41(10), 1553-1557 CODEN: TELEAY; ISSN: 0040-4039

PB Elsevier Science Ltd.

DT Journal

LA English

OS CASREACT 133:4952

GI

AB An efficient stepwise synthesis of oligoureas up to the nonamer, e.g. I, on solid support using O-succinimidyl-(9H-fluoren-9-ylmethoxycarbonylamino)ethylcarbamate derivs., e.g. II (R = PhCH2, Me), as activated monomers is described. These building blocks were readily prepd. starting from N-Fmoc-protected .beta.3-amino acids via Curtius rearrangement of the corresponding acyl azides and treatment of the resulting isocyanate with N-hydroxysuccinimide.

IT 270575-71-8P 270575-72-9P 270575-73-0P 270575-74-1P 270575-75-2P 270575-76-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(conversion of Fmoc-protected .beta.-amino acids to succinimidyl aminoethylcarbamate active monomers for prepn. of oligoureas)

RN 270575-71-8 CAPLUS

CN Carbamic acid, [(1S)-2-[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]-1-methylethyl]-, 9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 270575-72-9 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-2-methylpropyl]-, 9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RN 270575-73-0 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-3-methylbutyl]-, 9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 270575-74-1 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]-2-phenylethyl]-, 9H-fluoren-9-

ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 270575-75-2 CAPLUS

CN Carbamic acid, [(1S)-2-[4-(1;1-dimethylethoxy)phenyl]-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]ethyl]-, 9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 270575-76-3 CAPLUS

CN Carbamic acid, [(1S)-5-[[(1,1-dimethylethoxy)carbonyl]amino]-1-[[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]pentyl]-,
9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2002 ACS

AN 1999:769088 CAPLUS

DN 132:137681

TI Acyclic structural variants of growth hormone secretagogue L-692,429

AU Lin, Peter; Pisano, Judith M.; Schoen, William R.; Cheng, Kang; Chan, Wanda W.-S.; Butler, Bridget S.; Smith, Roy G.; Fisher, Michael H.; Wyvratt, Matthew J.

CS Department of Medicinal Chemistry, Rahway, NJ, 07065, USA

SO Bioorganic & Medicinal Chemistry Letters (1999), 9(22), 3237-3242 CODEN: BMCLE8; ISSN: 0960-894X

PB Elsevier Science Ltd.

DT Journal

LA English

GI

AB Starting with L-692,429 as a design template, several new acyclic growth hormone secretagogues were prepd. and evaluated for their hormone release activity in vitro. N-phenylamides derived by ring cleavage of L-692,429 were inactive. Arom. amino acid derivs. were active, the D-homophenylalanine derivs. being most active, with I having activity comparable to that of L-692,429.

IT 256479-80-8

RL: RCT (Reactant); RACT (Reactant or reagent)
(prepn. and activity of acyclic structural variants of growth hormone secretagogue L-692,429)

RN 256479-80-8 CAPLUS

CN Carbamic acid, [2-[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]-1,1-dimethylethyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

RE.CNT 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 6 OF 6 CAPLUS COPYRIGHT 2002 ACS

AN 1999:670476 CAPLUS

DN 132:78833

TI Effective preparation of O-succinimidyl-2~ (tert-butoxycarbonylamino)ethylcarbamate derivatives from .beta.-amino acids. Application to the synthesis of urea-containing pseudopeptides and oligoureas

AU Guichard, Gilles; Semetey, Vincent; Didierjean, Claude; Aubry, Andre; Briand, Jean-Paul; Rodriguez, Marc

CS Laboratoire de Chimie Immunologique, UPR 9021 CNRS Institut de Biologie Moleculaire et Cellulaire, Strasbourg, 67000, Fr.

SO Journal of Organic Chemistry (1999), 64(23), 8702-8705 CODEN: JOCEAH; ISSN: 0022-3263

PB American Chemical Society

DT Journal

LA English

GΙ

AB The authors report the application of Curtius rearrangement for the simple

conversion of N-Boc-protected .beta.-amino acids I [R = H, Me, Pr-i, CH2Ph, CH2CO2CH2Ph, CH(Me)OCH2Ph, (CH2)4NHCO2C6H4Cl-2] into the corresponding O-succinimidyl-2-(tert-butoxycarbonylamino)ethylcarbamate derivs. II. II are stable, cryst. products that react readily with amines

to form substituted ureas and then can be used as activated monomers in the synthesis of oligoureas.

IT 254100-96-4P 254100-97-5P 254100-98-6P 254100-99-7P 254101-00-3P 254101-01-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(synthesis of pseudopeptides and oligoureas from O-succinimidyl (Boc-amino)ethylcarbamate derivs., prepd. from .beta.-amino acids)

RN 254100-96-4 CAPLUS

CN Carbamic acid, [(1S)-2-[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]-

methylethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 254100-97-5 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-

pyrrolidinyl)oxy]carbonyl]amino]me

thyl]-2-methylpropyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 254100-98-6 CAPLUS

CN Carbamic acid, [(1S)-1-[[[[(2,5-dioxo-1-

pyrrolidinyl)oxy]carbonyl]amino]me

thyl]-2-phenylethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 254100-99-7 CAPLUS

CN Butanoic acid, 3-[[(1,1-dimethylethoxy)carbonyl]amino]-4-[[[(2,5-dioxo-

1 pyrrolidinyl)oxy]carbonyl]amino]-, phenylmethyl ester, (3S)- (9CI) (CA
 INDEX NAME)

Absolute stereochemistry. Rotation (-).

Absolute stereochemistry. Rotation (+).

RN 254101-01-4 CAPLUS

CN Carbamic acid, [(1S)-5-[[[(2-chlorophenyl)methoxy]carbonyl]amino]-1[[[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl]amino]methyl]pentyl]-,
1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RE.CNT 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

# L7 ANSWER 1 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8600960

Chemical Name (CN): <6-(2,5-dioxo-pyrrolidin-1-

yloxycarbonylamino)-5-(9H-fluoren-9ylmethoxycarbonylamino)-hexyl>-carbamic

acid

tert-butyl ester

Autonom Name (AUN): <6-(2,5-dioxo-pyrrolidin-1-

yloxycarbonylamino)-5-(9H-fluoren-9ylmethoxycarbonylamino)-hexyl>-carbamic

acid

tert-butyl ester

Molec. Formula (MF): C31 H38 N4 O8

Molecular Weight (MW): 594.66

Lawson Number (LN): 25671, 5573, 3111, 1762, 318

File Segment (FS): Stereo compound Compound Type (CTYPE): heterocyclic

Constitution ID (CONSID): 7289177
Tautomer ID (TAUTID): 8100536
Entry Date (DED): 2000/10/24
Update Date (DUPD): 2000/10/24

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \* Atom/Bond Notes:

1. CIP Descriptor: S

## Reference(s):

# L7 ANSWER 2 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8599397

Molec. Formula (MF): C33 H35 N3 O7

Molecular Weight (MW): 585.66

Lawson Number (LN): 25671, 14912, 5573, 1762, 318

File Segment (FS): Stereo compound Compound Type (CTYPE): heterocyclic

Constitution ID (CONSID): 7287362
Tautomer ID (TAUTID): 8098451
Entry Date (DED): 2000/10/24
Update Date (DUPD): 2000/10/24

### Reference(s):

 Guichard, Gilles; Semetey, Vincent; Rodriguez, Marc; Briand, Jean-Paul, Tetrahedron Lett., CODEN: TELEAY, 41(10), <2000>, 1553 - 1558; BABS-6233843

### Reference(s):

## L7 ANSWER 3 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8596110

Chemical Name (CN): <2-(9H-fluoren-9-ylmethoxycarbonylamino)-

3-

phenyl-propyl>-carbamic acid
2,5-dioxo-pyrrolidin-1-yl ester

Autonom Name (AUN): <2-(9H-fluoren-9-ylmethoxycarbonylamino)-

3-

phenyl-propyl>-carbamic acid
2,5-dioxo-pyrrolidin-1-yl ester

Molec. Formula (MF): C29 H27 N3 O6

Molecular Weight (MW): 513.55

Lawson Number (LN): 25671, 14535, 5573, 1762

File Segment (FS): Stereo compound Compound Type (CTYPE): heterocyclic

Constitution ID (CONSID): 7284645
Tautomer ID (TAUTID): 8096979
Entry Date (DED): 2000/10/24
Update Date (DUPD): 2000/10/24

## Reference(s):

## L7 ANSWER 4 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8593130

Chemical Name (CN): <2-(9H-fluoren-9-ylmethoxycarbonylamino)-

4-

methyl-pentyl>-carbamic acid
2,5-dioxo-pyrrolidin-1-yl ester

Autonom Name (AUN): <2-(9H-fluoren-9-ylmethoxycarbonylamino)-

2000/10/24

4 –

methyl-pentyl>-carbamic acid
2,5-dioxo-pyrrolidin-1-yl ester

Molec. Formula (MF): C26 H29 N3 O6

Molecular Weight (MW): 479.53

Update Date (DUPD):

Lawson Number (LN): 25671, 5573, 3048, 1762

File Segment (FS):

Compound Type (CTYPE):

Constitution ID (CONSID):

Tautomer ID (TAUTID):

Entry Date (DED):

Stereo compound
heterocyclic
7283153
8096101
2000/10/24

### Reference(s):

### L7 ANSWER 5 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8591827

Chemical Name (CN): <2-(9H-fluoren-9-ylmethoxycarbonylamino)-

3-

methyl-butyl>-carbamic acid
2,5-dioxo-pyrrolidin-1-yl ester

Autonom Name (AUN): <2-(9H-fluoren-9-ylmethoxycarbonylamino)-

3-

methyl-butyl>-carbamic acid
2,5-dioxo-pyrrolidin-1-yl ester

Molec. Formula (MF): C25 H27 N3 O6

Molecular Weight (MW): 465.50

Lawson Number (LN): 25671, 5573, 3047, 1762

File Segment (FS):

Compound Type (CTYPE):

Constitution ID (CONSID):

Tautomer ID (TAUTID):

Entry Date (DED):

Update Date (DUPD):

Stereo compound
heterocyclic
7281096
8095337
2000/10/24

## Reference(s):

#### L7 ANSWER 6 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8589843

Chemical Name (CN): <2-(9H-fluoren-9-ylmethoxycarbonylamino)-

propyl>-carbamic acid 2,5-dioxo-

pyrrolidin-1-

yl ester

Autonom Name (AUN): <2-(9H-fluoren-9-ylmethoxycarbonylamino)-

propyl>-carbamic acid 2,5-dioxo-

pyrrolidin-1-

yl ester C23 H23 N3 O6

Molec. Formula (MF): C23 H23 N3

Molecular Weight (MW): 437.45 Lawson Number (LN): 25671,

25671, 5573, 3028, 1762

File Segment (FS):
Compound Type (CTYPE):

Stereo compound heterocyclic

Compound Type (CTYPE):
Constitution ID (CONSID):
Tautomer ID (TAUTID):

7279295 8094868 2000/10/24

Entry Date (DED):
Update Date (DUPD):

2000/10/24 2000/10/24

## Reference(s):

# L7 ANSWER 7 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8459107

Chemical Name (CN): (S)-O-succinimidyl-2-(tert-

butoxycarbonylamino)-6-(2-

# chlorobenzyloxycarbonylamino)hexanoylcarbama

te

Autonom Name (AUN): <5-tert-butoxycarbonylamino-6-(2,5-dioxo-

2000/05/16

pyrrolidin-1-yloxycarbonylamino)-hexyl>-

carbamic acid 2-chloro-benzyl ester

Molec. Formula (MF): C24 H33 Cl N4 O8

Molecular Weight (MW): 541.00

Lawson Number (LN): 25671, 5229, 3111, 1762, 318

File Segment (FS):

Compound Type (CTYPE):

Constitution ID (CONSID):

Tautomer ID (TAUTID):

Entry Date (DED):

Stereo compound
heterocyclic
7175871
7974775
2000/05/16

Update Date (DUPD):

## Reference(s):

# L7 ANSWER 8 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8449483

Chemical Name (CN): (S)-O-succinimidyl-3-(benzyloxycarbonyl)-

2-

Autonom Name (AUN): (tert-butoxycarbonylamino)propylcarbamate 3-tert-butoxycarbonylamino-4-(2,5-dioxo-

pyrrolidin-1-yloxycarbonylamino)-butyric

acid benzyl ester

Molec. Formula (MF): C21 H27 N3 O8

Molecular Weight (MW): 449.46

Lawson Number (LN): 25671, 5228, 3398, 1762, 318

File Segment (FS): Stereo compound Compound Type (CTYPE): heterocyclic

Constitution ID (CONSID): 7167456
Tautomer ID (TAUTID): 7972673
Entry Date (DED): 2000/05/16
Update Date (DUPD): 2000/05/16

Reference(s):

#### ANSWER 9 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL L7

8447291 Beilstein Records (BRN):

Chemical Name (CN):

Molec. Formula (MF):

Lawson Number (LN):

File Segment (FS):

Entry Date (DED):

Update Date (DUPD):

Molecular Weight (MW):

Compound Type (CTYPE):

Tautomer ID (TAUTID):

Constitution ID (CONSID):

(2R, 3R)-O-succinimidyl-3-(benzyloxy)-2-

(tert-

butoxycarbonylamino)propylcarbamate Autonom Name (AUN):

<2-benzyloxy-1-<(2,5-dioxo-pyrrolidin-1-

yloxycarbonylamino)-methyl>-propyl>-

carbamic

acid tert-butyl ester

C21 H29 N3 O7

435.48

25671, 5228, 3142, 1762, 318

Stereo compound

heterocyclic

7165502

7968655 2000/05/16

2000/05/16

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ \end{array}$$

Reference(s):

### L7 ANSWER 10 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8441309

Chemical Name (CN): (S)-O-succinimidyl-2-(tert-

butoxycarbonylamino)-4-

phenylpropylcarbamate .

Update Date (DUPD):

Autonom Name (AUN): (2-tert-butoxycarbonylamino-3-phenyl-

2000/05/16

propyl)-

carbamic acid 2,5-dioxo-pyrrolidin-1-yl

ester

Molec. Formula (MF): C19 H25 N3 O6

Molecular Weight (MW): 391.42

Lawson Number (LN): 25671, 14535, 1762, 318

File Segment (FS): Stereo compound Compound Type (CTYPE): heterocyclic

Constitution ID (CONSID): 7162005
Tautomer ID (TAUTID): 7967470
Entry Date (DED): 2000/05/16

## Reference(s):

#### L7 ANSWER 11 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8435642

Chemical Name (CN): (S)-O-succinimidyl-2-(tert-

butoxycarbonylamino)-3-

methylpropylcarbamate

Molec. Formula (MF):

Autonom Name (AUN): <1-<(2,5-dioxo-pyrrolidin-1-

yloxycarbonylamino)-methyl>-2-methyl-

propyl>-

carbamic acid tert-butyl ester

C15 H25 N3 O6

Molecular Weight (MW): 343.38

Lawson Number (LN): 25671, 3047, 1762, 318

File Segment (FS): Stereo compound Compound Type (CTYPE): heterocyclic

Constitution ID (CONSID): 7155260
Tautomer ID (TAUTID): 7964272
Entry Date (DED): 2000/05/16
Update Date (DUPD): 2000/05/16

#### Reference(s):

### L7 ANSWER 12 OF 12 BEILSTEIN COPYRIGHT 2002 BEILSTEIN CDS MDL

Beilstein Records (BRN): 8430277

Chemical Name (CN): (S)-O-succinimidyl-2-(tert-

butoxycarbonylamino)propylcarbamate

Autonom Name (AUN): <2-(2,5-dioxo-pyrrolidin-1-

yloxycarbonylamino)-1-methyl-ethyl>-

#### carbamic

acid tert-butyl ester

Molec. Formula (MF): C13 H21 N3 O6

Molecular Weight (MW): 315.33

Lawson Number (LN): 25671, 3028, 1762, 318

File Segment (FS): Stereo compound Compound Type (CTYPE): heterocyclic

Constitution ID (CONSID): 7152522
Tautomer ID (TAUTID): 7963446
Entry Date (DED): 2000/05/16
Update Date (DUPD): 2000/05/16

### Reference(s):

L10 ANSWER 1 OF 3 MARPAT COPYRIGHT 2002 ACS

AN 132:64173 MARPAT

TI Preparation of labeling reactants for fluorescent labeling of biospecific

binding reactants

IN Takalo, Harri; Hovinen, Jari; Mukkala, Veli-matti; Liitti, Pivi; Mikola, Heikki

PA Wallac Oy, Finland

SO Eur. Pat. Appl., 26 pp.

CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

		PATENT NO.			KIND DATI		DATE	ATE			APPLICATION NO.				DATE				
	PI	EP	9672	05		A.	1	1999	1229		EF	199	99-66	50100	)	19990	0603		
			R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
				ΙE,	SI,	LT,	LV,	FI,	RO										
		US	6080	839		Α		2000	0627		US	199	98-10	04219	9	19980	0625		
	PRAI	US	1998	-1042	219	199	9806	25											
	OS	OS CASREACT 132:64173																	
	GI																		

$$\begin{bmatrix} G & R \\ A & A \\ N & R1 & R1 & I \end{bmatrix}$$

AB Novel pyridinediylbis (methylenenitrilo) tetrakisacetic acid labeling reactants, suitable for fluorescent labeling of biospecific binding reactants in solid-phase synthesis, were prepd. The novel labeling reactants (I) [wherein A = a bivalent arom. structure capable of absorbing

light or energy and transferring the excitation energy to a lanthanide ion

after the product made by solid-phase synthesis has been released from the  $\ensuremath{\mathsf{I}}$ 

used solid support, deprotected, and converted to a lanthanide chelate;  $\ensuremath{\mathtt{R}}$ 

= -Z(G1-NH-X)G2-E; X = a transient protecting group, e.g.
2-(4-nitrophenylsulfonyl)ethoxycarbonyl, trityl, 4-methoxytrityl,
4,4'-dimethoxytrityl, BOC, Fmoc; E = a carboxylic acid, its salt, active
ester (e.g. N-hydroxysuccinimido, nitrophenol, 2,4-dinitrophenol, or
pentafluorophenol), or halide; Z = the bridge point; G = a bridge
between

A and Z; G1 = a bridge between NH and Z; G2 = a bridge between E and Z; R1

= CO2R2; R2 = alkyl or (un)substituted Ph or benzyl] are particularly useful in the labeling of small mols. Thus, II was prepd. in a 4-step sequence involving (1) desilylation of Me (4-

trimethylsilylethynylphenoxy)

acetate (83%), (2) addn. to tetra(tert-Bu) 2,2',2'',2'''-[(4-bromopyridine-

2,6-diyl)bis(methylenenitrilo)]tetrakis(acetate) (75%), (3) deesterification of the phenoxyacetate with KOH (67%), and (4) amidation with .alpha.-Fmoc-lysine.HCl (56%). II was used for labeling of an estradiol deriv., incorporating four Eu(III) chelates, on a solid support

(no data).

### MSTR 1A

= 26

G6 = N

G3

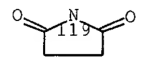
G8 = alkylene<(1-12)>

G10 = 81

G11 = 115

19<del>5 G</del>13

G13 = 119



DER: or salts MPL: claim 1

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L10 ANSWER 2 OF 3 MARPAT COPYRIGHT 2002 ACS

126:293352 MARPAT AN

Preparation of benzimidazoles for the prevention and/or the treatment of  $\mathtt{TI}$ bone diseases

Oku, Teruo; Kawai, Yoshio; Yatabe, Takumi; Sato, Shigeki; Yamazaki, IN Hitoshi; Kayakiri, Natsuko; Yoshihara, Kousei

Fujisawa Pharmaceutical Co., Ltd., Japan; Oku, Teruo; Kawai, Yoshio; PAYatabe, Takumi; Sato, Shigeki; Yamazaki, Hitoshi; Kayakiri, Natsuko; Yoshihara, Kousei

PCT Int. Appl., 146 pp. SO CODEN: PIXXD2

DTPatent

English LA

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE 19960905

PIWO 9710219 A1 19970320 WO 1996-JP2530

W: JP, US

RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT,

SE

EP 863881 A1 19980916 EP 1996-929540 19960905 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE,

FIJP 11513364

T2 JP 1996-511824 19991116 19960905

PRAI GB 1995-18552 19950911

WO 1996-JP2530 19960905

GΙ

---

ΑB The title compds. [I; R1 = acyl, (un) substituted lower alkenyl, lower alkyl; R2 = H, lower alkyl, lower alkoxy, etc.; R1R2 = lower alkylene, lower alkenylene (may include O, S, NH, N-alkyl); R3 = H, halo; R4 = (un) substituted heterocyclyl, aryl; A = CONR9, N(R10)CO (wherein R9, R10

H, (un) substituted lower alkyl)], and their pharmaceutically acceptable salts, inhibitors of bone resorption and bone metab., were prepd. Thus, hydrogenation of 1,2-dimethyl-4-nitro-1H-benzimidazole over 10% Pd/C in MeOH followed by reaction of the resulting 4-amino-1,2-dimethyl-1Hbenzimidazole with 2,6-dichlorobenzoyl chloride in the presence of Et3N

in ethylene chloride afforded I [R1, R2 = Me; R3 = H; R4 = 2,6-Cl2C6H3; A = NHCO]. Compds. I are effective at 0.1-1000 mg/body/day.

### MSTR 1

G17 = 61 / alkylamino<(1-6)> (SR (2) PO3H2)

DER: and pharmaceutically acceptable salts

MPL: claim 1

NTE: also incorporates claim 4

## MSTR 2

Ģ1—G2

G17 = 61 / alkylamino<(1-6)> (SR (2) PO3H2)

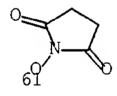
DER: and pharmaceutically acceptable salts

MPL: claim 1

# MSTR 3

Ģ1—G9

G17 = 61 / alkylamino<(1-6)> (SR (2) PO3H2)



DER: or reactive derivatives or salts

MPL: claim 4

```
L10
     ANSWER 3 OF 3 MARPAT COPYRIGHT 2002 ACS
     120:212035 MARPAT
AN
     Universal standard reagents for analyzing compounds having functional
TI
     groups, method of preparing same, and use thereof
     Patchornik, Avraham
IN
     Patchornik, Zipora, Israel
PA
     PCT Int. Appl., 45 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
LА
FAN.CNT 1
     PATENT NO.
                      KIND
                            DATE
                                            APPLICATION NO.
                                                             DATE
PΙ
     WO 9401771
                       Α1
                            19940120
                                            WO 1993-US6980
                                                             19930714
         W: AT, AU, BB, BG, BR, BY, CA, CH, CZ, DE, DK, ES, FI, GB, HU, JP,
             KP, KR, KZ, LK, LU, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD,
             SE, SK, UA, US, VN
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
     IL 102495
                            19980615
                       A1
                                            IL 1992-102495
                                                             19920714
     AU 9347844
                       Α1
                            19940131
                                            AU 1993-47844
                                                             19930714
     EP 650595
                       A1
                            19950503
                                            EP 1993-918367
                                                             19930714
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT,
SE
     JP 08505220
                       T2
                            19960604
                                            JP 1993-503596
                                                             19930714
                                           US 1995-362519
     US 5576216
                       Α
                            19961119
                                                             19950105
PRAI IL 1992-102495
                      19920714
     WO 1993-US6980
                      19930714
    A universal std. chem. reagent is described for quant. visual and
AB
     spectrometric anal. of compds. having reactive functional groups,
     including mixts. and homologs of the compds. The reagent comprises
     Q-B-f (Q = org. moiety which can be measured quant., visually by color,
```

compd.

spectroscopically, or fluorometrically; B = nonreactive org. bridging unit

linking Q to a reactive functional group f, the bridging unit being of sufficient length or size to prevent any possible interaction of Q that might alter its spectroscopic properties even upon derivatization; f = reactive group which can react with a compd. to form covalently bonded derivs.). Chlorodinitrobenzene was reacted with 3-aminopropanol in MeOH to make DNPNH(CH2)3OH (I). I enabled the prediction of the existence of self-catalytic reactions in acetylated glucose. DNPNH(CH2)3NHNH2 was

to analyze a triglyceride.

## MSTR 1C

used

G1---G2---G3

G1 = 93

$$G4 = NH$$

$$G6 = NH$$

$$G9 = 122$$

=> d ll; d his; log y
L1 HAS NO ANSWERS
L1 STR

Structure attributes must be viewed using STN Express query preparation.

(FILE 'HOME' ENTERED AT 15:15:32 ON 17 SEP 2002)

FILE 'REGISTRY' ENTERED AT 15:15:44 ON 17 SEP 2002

L1 STRUCTURE UPLOADED

L2 0 S L1

L3 14 S L1 FUL

FILE 'CAPLUS' ENTERED AT 15:16:20 ON 17 SEP 2002

L4 6 S L3

FILE 'BEILSTEIN' ENTERED AT 15:17:00 ON 17 SEP 2002

L5 0 S L1

L6 12 S L1 FUL

L7 12 S L6 NOT L4

FILE 'MARPAT' ENTERED AT 15:18:06 ON 17 SEP 2002

L8 0 S L1

L9 4 S L1 FUL

L10 3 S L9 NOT L4

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	111.49	621.43
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-1.77	-5.49

STN INTERNATIONAL LOGOFF AT 15:19:20 ON 17 SEP 2002